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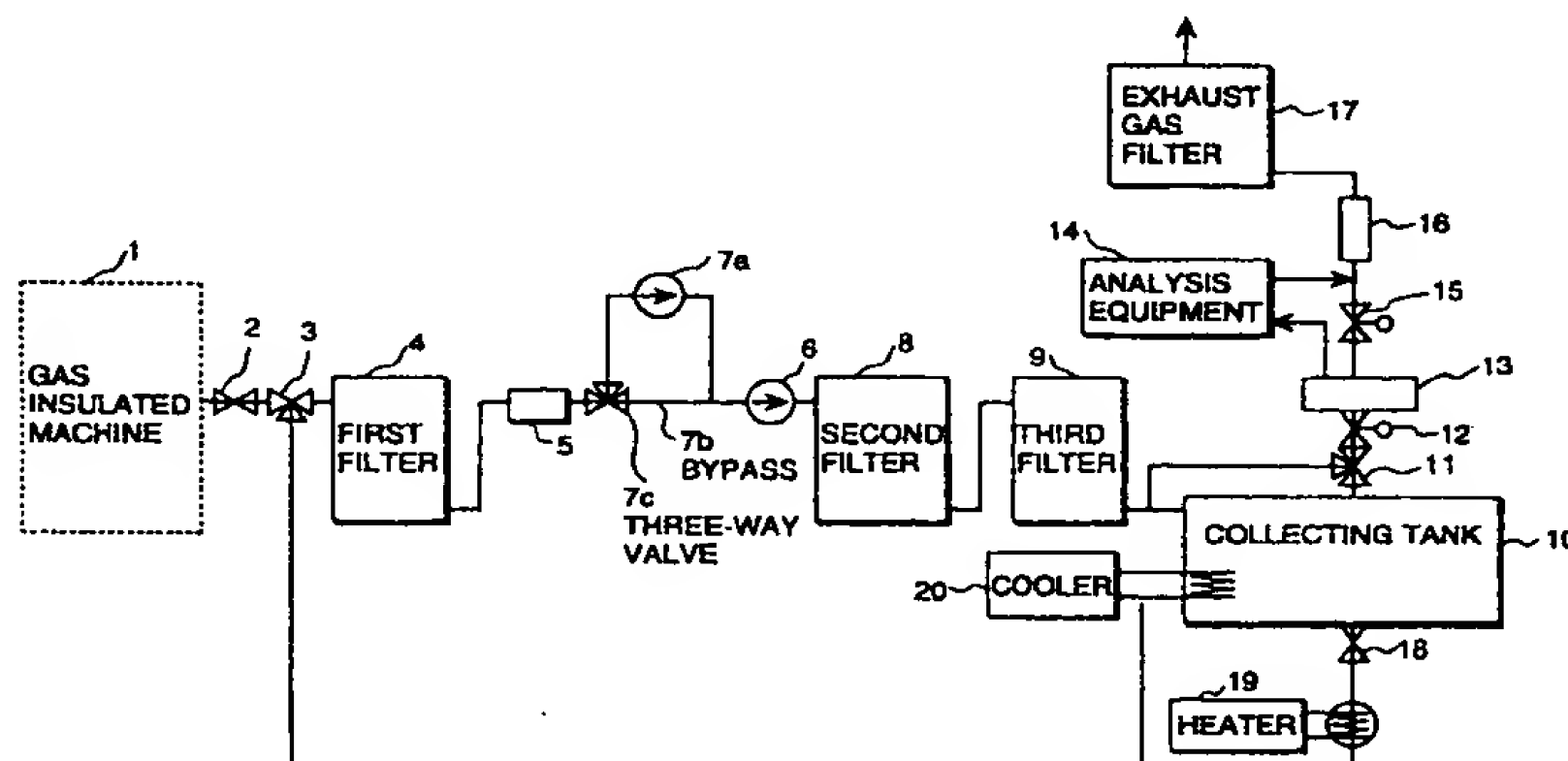
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(54) System and method for collecting and refining SF₆ gas

(57) SF₆ gas collected from the inside of a gas insulated machine (1) during maintenance and inspection is refined, and compositions of the refined SF₆ gas are analysed and confirmed to be reused at the site. Acidic gases are neutralised and removed by a dry method using filters (4, 8, 9) and the refined SF₆ gas is collected

in a collecting tank (10). The composition of the collected SF₆ gas after being refined is measured and confirmed by an analysis equipment (14) to quantitatively confirm whether or not the refined SF₆ gas is reusable.

FIG.1



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Description

Background of the Invention

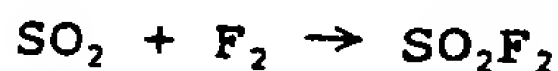
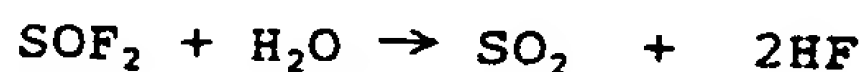
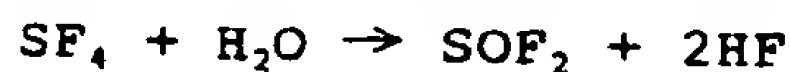
5 The present invention relates to a system for collecting and refining SF₆ gas at maintenance and inspection of an insulated machine using SF₆ gas as an insulating gas and a method therefor and, more particularly, to a system for collecting and refining SF₆ gas and a method therefor suitable for recovering a deteriorated SF₆ gas by collecting and refining.

10 Gas insulated machines commonly use SF₆ gas which is an inflammable insulating gas. For example, in the past transformers have employed an insulating oil as the insulating material, but an inflammable insulating gas is used instead of the insulating oil because the insulating oil has a probability of occurrence of fire or explosion in a case of trouble. SF₆ gas is widely used as the insulating gas. SF₆ gas has a global Warming property (hereinafter referred to as GWP) coefficient as large as approximately 25000 and accordingly has a large effect to the global Warming. Therefore, from the viewpoint of global warming prevention, it is preferable to prevent SF₆ gas, even a small quantity, from discharging to atmosphere. Further, the density of SF₆ gas is approximately five times as large as that of air, there is a possibility to cause an accident resulting in injury or death by an oxygen shortage when SF₆ gas is discharged inside a room because it stagnates in the bottom portion of the room or the lowermost portion of the building.

15 Furthermore, it is known that SF₆ gas is partly decomposed during using by arc discharge or the like, and part of the decomposed compositions react with moisture to produce acidic gases. Therefore, by-product refining gases are produced. That is, as shown in Table 1, SF₆ gas is partly decomposed by discharge or arc to produce acidic gases and metal fluorides such as HF, SOF₂, SO₂, H₂SO₃, SF₄, SF₂, S₂F₂ and so on.

Table 1

25 SF₆ → SF₄, SF₂, S₂F₂, F₂, Metal fluorides. When moisture exists, the following reactions occur.



40 In addition to these, there are some reports that CO₂ and so on are produced from an organic component such as winding portion, and that CF₄ is partially produced. As described above, although SF₆ gas is a useful gas as an insulating gas, SF₆ gas has a large global warming property coefficient and accordingly release of SF₆ gas to atmosphere is harmful to the global warming, and SF₆ gas may cause an accident such as an oxygen shortage when SF₆ gas is discharged inside a room because the density of SF₆ gas is large, and in addition to these SF₆ gas is harmful because it contains the acidic gases. Therefore, a method of collecting or recovering the deteriorated SF₆ gas is also required from social needs.

45 From the above reason, at maintenance and inspection of an insulated machine using SF₆ gas, a method of collecting and bringing back SF₆ gas and refining the SF₆ gas to be reused is proposed. For example, a decomposed gas treating system for a SF₆ gas insulated electric machine is disclosed in Japanese Patent Application Laid-Open No.9-856 proposes a method in which acidic gases among by-product gases contained in a deteriorated used SF₆ gas is removed by bubbling in an alkaline solution and then the SF₆ gas is filled in a compressed-gas cylinder to be brought back and refine the SF₆ gas.

50 Further, an article entitled "1512 An SF₆ gas collecting apparatus in a large electric power test site" presented at the National Meeting of the Institute of Electrical Engineers of Japan (1996) proposes a method in which SF₆ gas is refined to a reusable purity by (1) removing metal fluorides, (2) removing decomposed gases, (3) removing moisture and (4) removing impurities using an air removing block. This method is to remove acidic gases using an alkaline solution, and proposes a system which has a dew-point meter and an oxygen concentration meter so that quantities of moisture and air content can be monitored. This process can satisfy the SF₆ gas standard of "IEC-376 New Gas Stand-

ard" shown in Table 2.

Table 2

Moisture	lower than 15 wt ppm
Air	lower than 500 wt ppm
Hydrolytic fluoride (HF)	lower than 1.0 wt ppm

Although the social need that the deteriorated SF₆ gas is collected at inspection and maintenance and refined to be reused is increasing as described above, the technologies disclosed in Japanese Patent Application Laid-Open No.9-856 and described in the article entitled "1512 An SF₆ gas collecting apparatus in a large electric power test site" presented at the National Meeting of the Institute of Electrical Engineers of Japan (1996) are still technically insufficient to cope with the need. This is because the purity of the refined SF₆ gas is still not sufficiently confirmed and guaranteed, and the system for collecting and refining SF₆ gas is very large in size.

Initially, the confirmation and guarantee of gas purity will be described. SF₆ gas is an important factor to dominate the insulation performance of a gas insulated transformer or a gas insulated switch-gear. Therefore, in order to refine and reuse the SF₆ gas, it is required to confirm that the performance of the recovered SF₆ gas is equivalent to that of new SF₆ gas. If the recovered SF₆ gas contains impurities and the insulating performance is degraded, the apparatus is broken and damaged by occurrence of a short circuit inside the apparatus. Therefore, the collection and refinement requires confirmation by measuring the property of the recovered SF₆ gas. Table 3 shows an analysis result of an SF₆ gas used in a gas insulated switch-gear after repeating to open and close. In order to confirm that the performance of the recovered SF₆ gas is equivalent to that of new SF₆ gas, it is necessary to measure on the items and to confirm the concentrations. In order to do so, it is necessary to measure the items using measurement apparatuses matching to respective gases.

Table 3

Example of a result of analyzing components of a deteriorated SF ₆ gas			
No.	Item	Contents	Concentration (ppm)
1	Solids	Fluorides, Sulfides and others	
2	Acidic gases	SF ₄ , SF ₂ , S ₂ F ₂ , SOF ₂ , SO ₂ F ₂ , HF and the like	5000
3	Carbon dioxide	CO ₂	1500
4	PFC	CF ₄ , C ₂ F ₆ , C ₂ F ₄ and the like	1000
5	Air	O ₂ , N ₂	600
6	Moisture	H ₂ O	20

Further, since the gas insulated machine can be made small in size compared to that of the conventional method, the gas insulated machine in a substation in a town is usually installed in a basement form the viewpoint of effective use of the site area. In this case, since the gas insulating method has less fire potential compared to the conventional oil insulating method in addition to the advantage of small size, gas insulated machines are widely employed.

On the other hand, there are many substations of ground installed type in the suburb, the gas insulated machine capable of reducing the installation area is advantageous even in such a case.

The gas insulated machines are distributedly installed in many sites as described above, it is required that the system for collecting and refining the SF₆ gas to reuse it is small and portable. If the system for collecting and refining is not small and portable, it is required to fill the collected gas into a compressed gas-cylinder and to transfer it to a place where a refining apparatus is installed and then after refining to transfer it to the site again. Therefore, time period of the inspection process is lengthened and work for carrying in and out the compressed-gas cylinder is necessary. In a case where new SF₆ gas is filled after a deteriorated SF₆ gas is collected, delay in time period of the inspection process can be avoided, but work for carrying in and out of the compressed-gas cylinder of the collected gas and the compressed-gas cylinder of new SF₆ gas is required. In addition to this, the treatment of the collected deteriorated SF₆ gas is additionally required. In a case of a basement substation, the amount of work for carrying in and out of the collecting and refining system and of the compressed-gas cylinders becomes large and size of the transportation port also becomes

a limitation. Therefore, a small sized collecting and refining system is required.

Both of Japanese Patent Application Laid-Open No.9-856 and the article entitled "1512 An SF₆ gas collecting apparatus in a large electric power test site" presented at the National Meeting of the Institute of Electrical Engineers of Japan (1996) propose a wet method of refining deteriorated SF₆ gas where acidic gases are removed by letting the deteriorated SF₆ gas through lime-water solving calcium hydroxide in saturation. However, since the solubility of calcium hydroxide is small (0.16 g/100 g water at 20 °C) and accordingly the amount of the solution becomes large when it is used in an aqueous solution, there are problems in that the size of the apparatus becomes large and that a large amount of waste water is produced and a lot of time and effort are required in the transportation and treatment of the waste water. Particularly, since the waste water has a potential of leakage, the waste water needs to be carefully handled and managed, and there are various kinds of problems in handling the waste water in a limited place such as a basement.

Since the conventional SF₆ gas collecting system has only a gas collecting function but not a gas refining function, the conventional SF₆ gas collecting system is designed based on the viewpoint of how fast SF₆ gas is collected from a gas insulated machine in order to improve work efficiency. When SF₆ gas is filled in a gas insulated machine, for example, at a pressure of 5 kg/cm², the SF₆ gas is firstly collected into a collecting tank until pressure of the collecting tank and pressure of the gas insulated machine becomes equilibrium by opening a communicating pipe line, and then the remaining SF₆ gas is collected into the collecting tank using a compressor or a vacuum pump.

In a case of employing this method, pressure of the machine side is decreased from 5 kg/cm² to approximately 3 kg/cm² within a 1 to 2 minutes after opening the communicating pipe line. Then, the remaining SF₆ gas is collected using the compressor. However, approximately 1/3 of the gas has been collected by the opening of the communicating pipe line. Since an efficiency of the compressor decreases as the pressure in the machine side is decreased, an amount of collecting gas per unit time is decreased as the compressor is operated and the amount of collecting gas per unit time becomes the minimum at the end of the collecting work.

In a case where the processing rate of gas per unit time in the refining process is varied regardless of whether the process is of a wet method or a dry method, the refining process system needs to be designed so as to meet to the maximum processing gas flow rate. In a case where concentration of acidic gases in the SF₆ gas is reduced down to a predetermined value, design of the system of the wet method is performed by determining a used amount of alkaline solution to a unit volume of the treated gas (ratio of solution to gas) to a predetermined value. On the other hand, in a case of the dry method, design is also performed by determining a ratio of the treated gas volume treated in unit time to a volume of an adsorption removing agent (SV value). Therefore, the system needs to be designed so as to meet to the maximum gas flow rate when the processed gas rate varies, which leads to problems in that the system becomes large in size and consequently optimization and small-sizing of the system cannot be attained. Further, when the gas flow rate exceeds a value of design condition, a large amount of the alkaline solution is carried over by the gas in a case of the wet method to cause a trouble in the following process. Furthermore, regardless of the dry method or the wet method, there occurs a problem in that the initial removing performance cannot be attained.

Summary of the Invention

A first object of the present invention is to provide a system for collecting and refining SF₆ gas and a method of collecting and refining SF₆ gas which are small-sized and easy-operable in collection and refinement of SF₆ gas from an SF₆ gas insulated machine.

A second object of the present invention is to provide a system for collecting and refining SF₆ gas and a method of collecting and refining SF₆ gas which are capable of measuring and confirming compositions of SF₆ gas and capable of reusing the SF₆ gas in a short time at a working site of the SF₆ gas collecting and refining processing.

In order to attain the first object described above, a system for collecting and refining SF₆ gas in accordance with the present invention is characterized by comprising a first filter connected to a gas insulated machine; a constant flow rate supplier connected to a downstream side of the first filter; a compressor and a vacuum pump connected in parallel to a downstream side of the constant flow rate supplier; a collecting tank connected to a downstream side of the compressor; and a cooler for cooling the collecting tank, wherein the first filter is constructed by a filter filled with a dry adsorbent.

In addition to this, the system for collecting and refining SF₆ gas is characterized by further comprising a constant flow rate valve arranged in an inlet side of said compressor.

In order to attain the first object described above, a method of collecting and refining SF₆ gas in accordance with the present invention is characterized by comprising the steps of removing impurities from SF₆ gas from a gas insulated machine by averaging a flow rate of the SF₆ gas and letting the SF₆ gas flow through a first filter filled with a dry adsorbent; adsorbing and removing by-product gases by a second filter filled with a dry adsorbent; and collecting the SF₆ gas separated by the second filter by cooling and liquefying the SF₆ gas.

In order to attain the second object described above, a system for collecting and refining SF₆ gas in accordance

with the present invention is characterized by comprising at least a second filter arranged between the compressor and the collecting tank; and an analysis equipment including a gaschromatograph in a downstream side of the second filter.

In order to attain the second object described above, a method of collecting and refining SF_6 gas in accordance with the present invention is characterized by comprising the step of confirming compositions of the collected SF_6 gas by analyzing composition analysis items of moisture concentration, acidic gas concentration, carbon dioxide concentration, oxygen or nitrogen concentration and PFC concentration, and repeating the process of confirming said compositions and a process of returning the collected SF_6 gas to the first filter until the compositions become preset values.

Brief Description of the Drawings

These and other objects, features and advantages of the present invention will be understood more clearly from the following detailed description with reference to the accompanying drawings, wherein

FIG. 1 is a diagram showing the construction of a first embodiment of a system for collecting and refining SF_6 gas in accordance with the present invention;

FIG. 2 is a graph showing the relationship between collecting processing time and compressor inlet pressure;

FIG. 3 is a graph showing the relationship between collecting processing time and collecting and processing rate of insulating gas;

FIG. 4 is a diagram showing the construction of a second embodiment of a system for collecting and refining SF_6 gas in accordance with the present invention;

FIG. 5 is a graph showing change of concentration of processed gas with operation of a neutralizing and removing agent;

FIG. 6 is a diagram showing the construction of a third embodiment of a system for collecting and refining SF_6 gas in accordance with the present invention;

FIG. 7 is a diagram showing the construction of a fourth embodiment of a system for collecting and refining SF_6 gas using a film filter in accordance with the present invention; and

FIG. 8 is a diagram showing the construction of a modification of the embodiment of FIG. 7 of a system for collecting and refining SF_6 gas using a film filter.

Description of the Preferred Embodiments

A first embodiment of the present invention will be described below, referring to FIG. 1 to FIG. 3. FIG. 1 is a diagram showing the construction of a first embodiment of the system for collecting and refining SF_6 gas. FIG. 2 is a graph showing the relationship between collecting processing time and compressor inlet pressure. FIG. 3 is a graph showing the relationship between collecting processing time and collecting and processing rate of insulating gas.

The system for collecting and refining SF_6 gas of this embodiment is constructed as follows. As shown in FIG. 1, a gas insulated machine 1 is connected to a first filter 4 through a dividing valve 2 and a dividing three-way valve 3. An insulating gas, SF_6 , of 10 m^3 at 5 kg/cm^2 is filled in the gas insulated machine 1. The first filter 4 is connected to a second filter 8 through a constant flow rate supplier 5 and a compressor 6. A vacuum pump 7a and a by-pass pipe 7b are arranged between the constant flow rate supplier 5 and the compressor 6 in parallel, and a flow path of the vacuum pump 7a or the by-pass pipe 7b is selected through a three-way valve 7c. The three-way valve 7c connects the constant flow rate supplier 5 and the compressor 6 through the by-pass pipe 7b in the initial stage of operation, and connects the constant flow rate supplier 5 and the compressor 6 through the vacuum pump 7a when the pressure in the gas insulated machine 1 is decreased. Here, a pressure reducing valve is used as the constant flow rate supplier 5. The second filter 8 is connected to a third filter 9. The third filter 9 is connected to a collecting tank 10 having a cooler 20, and the collecting tank 10 is connected to a sampling header 13 through a sampling position switching valve 11 and a pressure reducing valve 12. The sampling header 13 is connected to an analysis equipment 14 and also connected to a flow meter 16 through an exhaust gas valve 15. Further, the analysis equipment 14 and the flow meter 16 are connected to each other in the downstream of the exhaust gas valve 15. The flow meter 16 is connected to an exhaust filter 17. On the other hand, the collecting tank 10 is connected to the dividing three-way valve 3 through a discharge valve 18 and a heater 19, and a branch portion is arranged between the heater 19 and the three-way valve 3, and the branch portion is connected to the sampling position switching valve 11.

Operation of the system for collecting and refining SF_6 gas constructed as described above will be described below. A deteriorated insulating gas in the gas insulated machine 1 is supplied to the first filter through the three-way valve by opening the dividing valve 2. The pressure reducing valve as the constant flow rate supplier 5 is fully closed. When the dividing valve 2 is opened and the inlet pressure at the constant flow rate supplier 5 becomes nearly equal to the pressure in the gas insulated machine, the pressure reducing valve is gradually opened and controlled so that the outlet pressure of the pressure reducing valve becomes 1.5 kg/cm^2 . Under this state, the compressor 6 is started to operate

to start collecting of the SF_6 gas. Since the pressure reducing valve is controlled so as to maintain the inlet pressure of the compressor to 1.5 kg/cm^2 , the collecting capacity of the compressor 6 is maintained at a constant value. Therefore, the collecting and processing rate of the SF_6 gas can be maintained to a constant value. Thereby, required amounts of adsorbent, neutralizer and the like can be minimized to necessary and minimum values. For example, the collecting amount of the SF_6 gas per unit time in the initial stage is set to $1 \text{ N m}^3/\text{min}$. Further, a fluid resistor such as an orifice or the like may be used as the constant flow rate supplier 5.

Since among impurities contained in the gas it is necessary to separate acidic gases and solid components such as fluorides and sulfides which are ill affect to machines, the neutralizing and removing agent is arranged inside the first filter 4 at a place in the upstream side of the compressor 6 and the vacuum pump 7. Since the first filter 4 is depressurized to a negative pressure of 10 Torr in the last stage, a neutralizing and removing agent of a physically adsorbing type is not preferable but a neutralizing and removing agent of a calcium group alkali base is preferable. The reason is that the neutralizing and removing agent of a physically adsorbing type has a characteristic of discharging substances once having been adsorbed under a depressurized condition, and accordingly activated carbon is not suitable. Therefore, in this embodiment, a granular neutralizing and removing agent having main components of CaO and Ca(OH)_2 is used. The neutralizing and removing agent fixes the acidic gases as calcium salts, and water produced by the fixing reaction is absorbed in CaO and carbon dioxide is also fixed and removed as carbonic salts by CaO and Ca(OH)_2 . A quantity of the neutralizing and removing agent to a quantity of the gas to be processed is set to $\text{SV} = 500 \text{ h}^{-1}$. Since the processing rate of the gas is $1 \text{ N m}^3/\text{min}$ ($60 \text{ N m}^3/\text{h}$), the quantity of the neutralizing and removing agent becomes approximately 120 liters. By doing so, the concentration of acidic gas after processed can be reduced below 1 ppm. Further, the solid components of fluorides and chlorides are also filtered and removed by the granular adsorbent layer.

An adsorbent selectively adsorbing a strong acidic gas of HF gas is contained in the first filter 4 to adsorb HF gas while the insulating medium is passing through the vessel of the first filter and to neutralize the insulating medium as a result. The method of neutralizing can be employed is a general neutralizing method of allowing the gas to pass through an alkaline solution. A dry neutralizing method of using an adsorbent can improve handling capability since no alkaline solution is used.

By using the granular neutralizing and removing agent having main components of CaO and Ca(OH)_2 to the first filter, the acidic gas after-processed can be removed and the water produced in an approximately equivalent quantity of the acidic gases accompanied by removing the acidic gases can be also removed. Further, carbon dioxide and solid components can be also removed and separated. Since the removal is performed by chemically changing the acidic gases and the like to the other substances to fix and separate, the removed substances cannot be released again even if the pressure of the first filter is depressurized down to near a vacuum during the collecting operation. In addition to these, since the components and the pipes in the system line up to the first filter 4 is exposed to the acidic gases, the components and pipes are protected by nickel plating (Kanigin plating) or the like.

The gas removed the acidic gases by the first filter 4 is supplied to the second filter 8 through the compressor 6. An adsorbent composed of activated carbon or zeolite as the main component is used for the second filter 8. The second filter 8 is arranged for backing up the first filter 4 and for adsorbing and separating by-product gases other than the acidic gases. The second filter 8 cannot be not brought in a negative pressure condition since it is arranged in the downstream side of the compressor 6 and the vacuum pump 7, and accordingly the substances once adsorbed and separated cannot be released again. Since an activity of adsorbing reaction of the activated carbon or the zeolite is high, the amount of the adsorbent of the second filter 8 is 60 liters by setting $\text{SV} = 1000 \text{ h}^{-1}$.

The gas of which the by-product gases have been adsorbed and separated by the second filter is supplied to the third filter 9. In the third filter 9, a molecular sieve is used for removing moisture, and the moisture concentration can be reduced to nearly zero by the back-up of the first filter 4. Since the absorbing characteristic of the molecular sieve is excellent, the amount of the molecular sieve is 30 liters by setting $\text{SV} = 2000 \text{ h}^{-1}$.

The deteriorated SF_6 gas processed by the individual filters as described above is collected and stored in the collecting tank 10 by being cooled and liquefied using the cooler 20. The SF_6 gas in the collecting tank 10 is separated into gas phase and a liquid phase, and a part of the SF_6 gas in gas phase in the upper portion of the collecting tank 10 is supplied to the analysis equipment 14 through the pressure reducing valve 12 and the sampling header 13 to perform composition analysis of the collected gas. Since the SF_6 gas is cooled and liquefied, non-condensable gases are gathered to the SF_6 gas in gas phase in the upper portion of the collecting tank 10 to increase the impurity concentration. The analysis equipment 14 comprises an acidic gas detector, a gas chromatograph, an oxygen meter and a dew point meter. The concentration of the acidic gases is measured by the acidic gas detector, and the concentrations of the carbon dioxide and the PFC gas are measured by the gas chromatograph. Further, the concentration of the air is measured by the oxygen meter, and the concentration of the moisture is measured by the dew point meter.

The supply of gas to the analysis equipment 14 is continuously performed, and accordingly the latest gas composition of the collected gas is analyzed and confirmed. The gas after completed the analysis is discharged to the external through the exhaust gas filter 17 which is filled with activated carbon or the like. The SF_6 gas is adsorbed in the exhaust gas filter 17 to prevent the SF_6 gas having a large global warming property coefficient from leaking to outside the sys-

tem. Otherwise, instead of using the exhaust gas filter 17, the exhaust gas may be collected into a vinyl bag or the like and the collected gas may be burned in an incinerator since the quantity of the exhaust gas is small.

If it is judged from the analysis result of the gas compositions by the analyzing equipment 14 that the purity of the gas does not reach the target value, the discharge valve 18 of the collecting tank 10 is opened, and the liquefied gas is heated with the heater 19 to be vaporized and returned to the first filter 4 through the dividing three-way valve 3.

As described above, the process of confirming the compositions and the process of returning the collected SF₆ gas to the first filter until the compositions become the target values.

Although air among the impurities leaks and mixes into the system, it is difficult to separate some of non-condensable gases such as air by adsorbing and removing. Therefore, if a result of the analysis shows that the concentration of air is high, the impurities such as air is selectively exhausted by opening the exhaust gas valve 15 to exhaust the gas in the gas phase of the collecting tank while the flow rate is being confirmed using the flow meter 16 because air (O₂, N₂) is a non-condensable gas having a lower condensing temperature compared to SF₆ gas and exists in the gas phase. PFC gas and carbon dioxide having lower condensing temperatures compared to SF₆ gas can be also selectively exhausted to the external in the similar manner. In that time, the SF₆ gas is prevented from being discharged outside the system by the exhaust gas filter 17. Further, similar to the above, the exhaust gas may be collected into a vinyl bag or the like and the collected gas may be burned in an incinerator since the quantity of the exhaust gas is small.

As the collection of the SF₆ gas is progressed in the manner as described above, pressure in the gas insulated machine 1 is gradually decreased and becomes lower than a set value of the pressure reducing valve as the constant flow rate supplier 5. Although the pressure reducing valve is in the fully opened state at that time, the collecting rate is decreased lower than the preset value and cannot exceed the design flow rate of the filter because the efficiency of the blower is decreased due to lowering of the pressure. The collection of the SF₆ gas is further progressed, and at the time when the pressure reaches approximately -0.1 to -0.2 kg/cm², the vacuum pump 7 is started to operate and the SF₆ gas is collected using the both machines. Finally, the SF₆ gas is collected as much as possible by reducing the pressure inside the gas insulated machine 1 up to 10 Torrs. By doing so, the amount of the remaining SF₆ gas inside the gas insulated machine 1 can be reduced and accordingly the amount of discharged SF₆ gas can be reduced. At the time of completion of collection, the final analysis of composition of the collected gas is performed to confirm that the composition is within the range of the required condition. Then, the gas insulated machine is opened to perform inspection and maintenance of the inside portions. After completion of the inspection and maintenance, the inside of the gas insulated machine 1 is evacuated to 10 Torrs to exhaust the air inside the gas insulated machine 1. Then, the collected and refined SF₆ gas is refilled in the gas insulated machine 1.

The refilling is performed by opening the exhaust valve 18 and heating and gasifying the cooled and liquefied SF₆ gas using the heater 19 to supply the SF₆ gas to the gas insulated machine 1. At that time, the composition of the refilled gas is analyzed and confirmed by switching the sampling position switching valve 11.

The SF₆ gas gasified by the heater 19 is supplied to the gas insulated machine 1 through the dividing three-way valve 3 and the dividing valve 3. In a case where the analysis result of the gas composition after collection and refining of the SF₆ gas shows that the purity of the gas is not sufficient, similar to the case of refilling after opening the exhaust valve 18 and heating and gasifying the cooled and liquefied SF₆ gas using the heater 19, the SF₆ gas is supplied to the first filter 4 through the dividing three-way valve 3 to repeat the same process as the process at collecting and refining time.

New SF₆ gas is additionally supplied from a compressed gas cylinder by the amount of the SF₆ gas consumed by being decomposed inside the gas insulated machine 1 and the amount of the SF₆ gas consumed by the analysis and the like so that a necessary amount of SF₆ gas is filled in the gas insulated machine. Further, the SF₆ gas adsorbed to the activated carbon in the exhaust gas filter 17 is burning-treated together with the activated carbon and not discharged outside the system.

By removing and separating the by-produced impurity gases contained in the deteriorated SF₆ gas not by wet but by the dry processing as described above, the system for collecting and refining SF₆ gas of this embodiment can be made small not only in the size of the adsorbing and removing components but also in the size of the total system. For example, when calcium hydroxide of 100 g is used as the saturated aqueous solution, the volume becomes approximately 63 liters because the solubility is small. When calcium hydroxide of 100 g is used as a 5 % concentration aqueous solution, the volume becomes approximately 2 liters. However, it is necessary to stir the solution uniformly. On the other hand, when calcium hydroxide is formed in granular shape or particle shape and used by dry method, volume of calcium hydroxide of 100 g is approximately 0.14 liter since the bulky density is approximately 0.7 g/cc. Even if the available efficiency is assumed to be nearly 70 % because of dry method, the volume becomes only approximately 0.2 liter. By treating the gas by the dry method as described above, the volume of the system can be reduced below 1/10. Further, the amount of the secondary waste can be also reduced below 1/10, and can be easily handled because no liquid waste is produced.

In addition to these, it is preferable from the viewpoint of work efficiency that the work of collecting the SF₆ gas from the gas insulated machine is performed at the installation site of the gas insulated machine, and after completion of

inspection the refined SF_6 gas collected from the gas insulated machine is refilled to the gas insulated machine. However, it is generally necessary that the measurement of the collected SF_6 gas is performed by sending the samples to a factory or a place where an analysis equipment is placed to be measured. Since the work needs to be temporary stopped in such a case, new SF_6 gas is often filled in the gas insulated machine. This is because the gas insulated machine is an electric supply infrastructure apparatus which is required to be recovered in a short time.

The system for collecting and refining SF_6 gas of this embodiment has an equipment capable of quantitatively analyzing and measuring the compositions of the collected and refined SF_6 gas in place, and accordingly it is possible to judge at the collecting site whether or not the SF_6 gas is reusable and to smoothly perform reuse of the SF_6 gas. That is, the measurement of the gas can be performed by selecting a measurement apparatus for each of the gases as shown in Table 4. By combining the analysis equipment for gas composition with the system for collecting and refining SF_6 gas and by measuring and confirming the composition of the gas after collecting and refining in place in a short time, as described above, after refining the collected SF_6 gas the refined gas can be speedy and effectively reused.

Table 4

Examples of applicable measurement apparatus			
No.	Item	Measurement Apparatus	Measurable Lower Limit (ppm)
1	Acidic gases	Acidic gas detector, Gas chromatograph	approximately 0.5
2	Carbon dioxide	Gas chromatograph	approximately 0.5
3	PFC	Gas chromatograph	approximately 10
4	Air	Oxygen meter	approximately 10
5	Humidity	Dew point meter, Humidity meter	approximately 5

A key point of improving work efficiency in the conventional system is how to collect the SF_6 gas in a short time, and consideration is paid only on collecting the SF_6 gas, but not on averaging the collecting flow rate. However, in order to reuse the SF_6 gas, how to refine the SF_6 gas to a high purity is an important factor to improve the work efficiency. If the purity is low, the SF_6 gas cannot be reused or the SF_6 gas needs to be refined again.

The maximum flow rate during collecting period occurs in the initial period of the collecting work. That is, the maximum flow rate during collecting period occurs in a period of 1 to 2 minutes when the pressures in the gas insulated machine and the collecting tank become in balance after connecting the gas insulated machine filled with SF_6 gas to the collecting tank with a communicating pipe and opening the communicating pipe, and in the following short period when the compressor for collecting the gas is started to operate and the pressure inside the gas insulated machine is reduced to a certain value. Since the compressor more efficiently collects the gas as the pressure in the suction side is higher, the efficiency is decreased as the pressure in the suction side decreases and consequently the collecting flow rate is reduced as the pressure in the suction side decreases. The maximum flow rate occurs in the period of opening the communicating pipe and the following short period after starting the compressor. Therefore, by performing constant flow rate control of the collecting flow rate during those periods, the collecting time is lengthened by 10 to 20 % compared to that in the conventional system. However, the collecting flow rate can be made constant and averaged, the processing system can be made small in size, and the refining processing can be improved.

In the past, collection of the SF_6 gas has been performed in a manner as shown by the broken line in FIG. 5. That is, collection of the SF_6 gas is started when the inlet pressure of the compressor of the collecting system is a value equal to the pressure in the gas insulated machine, and the inlet pressure is rapidly decreased as shown in FIG. 2 as the collection of the SF_6 gas is progressed. During the period, the collecting flow rate changes along the curve shown in FIG. 3 which is similar to the pressure change shown in FIG. 2. However, the maximum process capacity of the refining apparatus needs to have a process capacity meeting to the point A.

On the other hand, according to the present invention, since the pressure reducing valve 5 is arranged in the inlet of the compressor 6 of the collecting system and the SF_6 gas is collected with keeping the inlet pressure of the compressor 6 and controlling the collecting flow rate to a constant flow rate, the collecting characteristic becomes as shown by the solid line in FIG. 5. Since the inlet pressure of the compressor is kept constant at the value of the point B, the collecting flow rate is also kept constant. When the pressure in the gas insulated machine side is decreased lower than a preset value of the pressure reducing valve, the pressure reducing valve is fully opened at the point D and the inlet pressure of the compressor is decreased to decrease the collecting flow rate. By doing the constant flow rate control of the collecting flow rate, the time period necessary for collecting and refining the SF_6 gas is lengthened by Δt , but it is

lengthened only by 10 to 20 % compared to the total time period. By doing so, the process capacity of the system for collecting and refining can be reduced from the point A to the point C. Thereby, the necessary process capacity can be reduced to 1/3 to 1/4.

A second embodiment of the present invention will be described, referring to FIG. 4 and FIG. 5. FIG. 4 is a diagram showing the construction of this embodiment of a system for collecting and refining SF₆ gas. FIG. 5 is a graph showing change of concentration of processed gas with operation of a neutralizing and removing agent.

The system for collecting and refining SF₆ gas of this embodiment is constructed similarly to the embodiment shown in FIG. 1. However, in this embodiment, a four-way switching valve 21 and a four-way switching valve 22 are arranged in the downstream side of the dividing three-way valve 3, and two first filters 4a, 4b are arranged in parallel through the four-way switching valve 21 and the four-way switching valve 22. The two first filters 4a, 4b are connected double in parallel as shown in FIG. 4 to improve the availability of the neutralizing and removing agent.

It is confirmed that the concentration of the acidic gases is the highest among the impurities in the deteriorated SF₆ gas from an inspection and analysis result of compositions of the deteriorated SF₆ gas. In a case of adsorption or reaction fixing, an effective amount of the adsorbent or the neutralizing and removing agent is decreased with operation and the adsorbent needs to be exchanged because of reducing of the removing efficiency. It can be understood from FIG. 5 showing change of concentration of the processed gas with operation of the neutralizing and removing agent that when processing is performed by filling a designed amount of the neutralizing and removing agent, the concentration of the acidic gases is gradually decreased in the inside of the filter and at the outlet the acidic acid can be removed below a desired concentration, as shown by the solid line. However, when the neutralizing and removing agent in the inlet portion of the filter is gradually consumed with operation, the concentration of the acidic gases inside the filter changes as shown by the broken line and the filled neutralizing and removing agent needs to be exchanged in order that the concentration of the acidic gases at the outlet attains the reference concentration. In this case, although the amount of the consumed neutralizing and removing agent is far less than a half of the total amount, the whole amount of the filled agent needs to be exchanged because the initial performance as the filter cannot be displayed.

Therefore, in this embodiment, two sets of the first filters filled with Ca group alkali having the largest load are used by arranging them in series as shown in FIG. 4. The collected SF₆ gas is initially supplied to the first filter 4a by switching the flow path using the four-way switching valve 21. The deteriorated SF₆ gas having passed through the first filter 4a is supplied to the first filter 4b through the four-way switching valve 22 and the four-way switching valve 21 to remove the acidic gases.

When the concentration of the acidic gas exceeds the reference value by monitoring the outlet concentration of the acidic gas using an acidic gas monitor 23, the neutralizing and removing agent filled in the first filter 4a is exchanged and the flow path is changed so that the deteriorated SF₆ gas flows through in order of the four-way switching valve 21, the first filter 4b, the four-way switching valve 22, the four-way switching valve 21, the first filter 4a, the four-way switching valve 22 and the compressor 6. By operating in such a manner, the total amount of the neutralizing and removing agent in each of the filters can be consumed out, and consequently the amount of the neutralizing and removing agent used can be reduced to approximately 1/2 compared to in a case of operating one set of the filter. As a result, in addition of the reduction in the procurement cost of the neutralizing and removing agent, the disposal cost of the secondary waste can be reduced to 1/2 and the system can be made smaller since the availability of the neutralizing and removing agent can be increased twice.

A third embodiment of the present invention will be described, referring to FIG. 6. FIG. 6 is a diagram showing the construction of this embodiment of a system for collecting and refining SF₆ gas.

As shown in FIG. 6, the collecting and refining system of this embodiment is constructed by combining an existing SF₆ gas collecting system 24 and an existing SF₆ gas refining system 25. In order to collect the SF₆ gas from the gas insulated machine, the SF₆ gas collecting system 24 is used. The existing SF₆ gas collecting system 24 is designed not in taking the refining process into consideration, but has only a function to collect the SF₆ gas from the gas insulated machine 1. By adding a gas refining function to this system, the existing system can be effectively used as a series of systems for collecting and refining SF₆ gas. The added SF₆ gas refining system 25 is mainly composed of filters and a composition analysis equipment 14 for collected gas.

The SF₆ gas collecting system 24 collects the SF₆ gas from the gas insulated machine 1 to the collecting tank 10 through the dividing valve 2 and the dividing three-way valve 3 using the compressor 6 and the vacuum pump 7, and the collected SF₆ gas is cooled in the collecting tank 10 by the cooler 20 to be liquefied. When the gas is refilled, the SF₆ gas is supplied to the gas insulated machine 1 through the dividing three-way valve 3 and the dividing valve 2 by opening the discharge valve 18 of the collecting tank 10 and heating using the heater 19 to evaporate the SF₆ gas.

When the SF₆ gas collecting system 24 and the SF₆ gas refining system 25 are combined, the system structure and the operation become as follows. The SF₆ gas in the gas insulated machine 1 is supplied to the first filter 4 through the dividing valve 2 and the dividing three-way valve 3. The dividing three-way valve 3 and the first filter 4 is connected by a pressure resistant and vacuum proof hose or pipe. The SF₆ gas passed through the first filter 4 is supplied to the compressor 6 or the vacuum pump 7 of the SF₆ gas collecting system 24 from the SF₆ gas refining system 25 through

the constant flow rate supplier 5. The SF₆ gas compressed by the compressor 6 or the vacuum pump 7 is supplied to a second filter 8 and a third filter 9 to remove impurities and then stored in the collecting tank 10. In the collecting tank 10, the SF₆ gas is cooled to be liquefied.

By separating into the collecting system and the refining system as the SF₆ gas collecting system 24 and the SF₆ gas refining system 25, each system can be made small and made better in portability. Further, when there is an existing collecting system, a series of collecting SF₆ gas and refining SF₆ gas can be performed by adding a new SF₆ gas refining system 25.

A fourth embodiment of the present invention will be described, referring to FIG. 7 and FIG. 8. Each of FIG. 7 and FIG. 8 is a diagram showing the construction of this embodiment of a system for collecting and refining SF₆ gas.

The system for collecting and refining SF₆ gas of this embodiment is constructed similar to the embodiment shown in FIG. 1. However, in the system for collecting and refining SF₆ gas of this embodiment, a fourth filter 30 is arranged between the third filter 9 and the collecting tank 10. In the fourth filter 30, a porous hollow thread film for selectively separating air is installed.

A produced amount, that is, a concentration of gas component composition produced in the deteriorated SF₆ gas in which SF₆ gas is deteriorated is seldom deviated largely from the values shown in Table 3. However, air enters into the system by in-leakage from the outside of the system, and the concentration of air possibly becomes 100 times or 1000 times of the concentration shown in Table 1 if even one of fastened positions is loosed. Further, in order to reduce an amount of SF₆ gas discharged to atmosphere, before releasing the gas insulated machine, the inside of the system other than the gas insulated machine is brought to a considerable vacuum condition in order to evacuation-collect SF₆ gas as much as possible. In this condition, potential of air to leak into the system from outside the system is increased.

On the other hand, air containing N₂, O₂ can be adsorbed and removed to a certain degree by an adsorbent of active carbon group, but the amount is as small as approximately 1 % of the amount of the adsorbent. Therefore, the air soon breaks through the adsorbent. The air may be discharged out of the system from the collecting tank 10 as a non-condensable gas. However, this case is not preferable because the SF₆ gas is also released outside the system together with the air. Therefore, it is necessary to selectively separate and remove only the air among the collected gases.

This embodiment realizes the selective separation and removal of the air using a porous hollow thread film having selective property. The acidic gases and the carbon dioxide gas accompanied by the SF₆ gas are removed by the first filter 4, and the by-product gases are removed in the second filter 8. Further, the moisture is removed in the third filter 9. The air containing N₂, O₂ is separated and removed by the porous hollow thread film filter in the fourth filter 30. This makes use of that the molecular diameters of N₂, O₂ gases are smaller than the molecular diameters of SF₆ gas, and only N₂, O₂ gases are separated and released outside the system using the film having a hole diameter through which N₂, O₂ gases pass but SF₆ gas does not pass. The gas is pumped into the fourth filter 30 by the compressor 6. Therefore, it is sufficient to place the film having the required property and no driving source for pressurizing the supplied gas is necessary because a positive pressure acts in the forth filter. Different from the filter of filling agent type, the film filter can cope with a large amount of air in-leaking because the film filter cannot be broken through by the air, and the film filter does not produce the secondary waste such as a disposed filling agent (disposed adsorbent).

In the embodiment shown in FIG. 8, the fourth filter 30 is arranged between the collecting tank 10 and the inlet of the compressor 6, and a branch portion is arranged in a position of the upstream side of the compressor 6 and the downstream side of the vacuum pump 7, and the forth filter 30 is connected to the branch portion through a pressure reducing valve 31. The fourth filter 30 is further connected to the analysis equipment 14, and the porous hollow thread film is arranged for selectively separate air is installed in the fourth filter 30.

In the embodiment shown in FIG. 8, the gas passes through the fourth filter 30 only once. In the system of FIG. 8, a predetermined amount of the gas is re-circulated by supplying the gas from the gas phase portion of the collecting tank 10 where the concentration of air of a non-condensable gas is high to the inlet portion of the compressor 6 through the pressure reducing valve 31. The re-circulation flow rate is controlled by the pressure reducing valve 31. Since the collecting tank 10 in this case is in a positive pressure, no driving source for applying a positive pressure to the film is necessary.

As described above, the efficiency of separating and removing the air can be substantially improved by re-circulating the gas in the gas phase portion where the concentration of air of a non-condensable gas is high in the collecting tank 10 and processing the gas using the fourth filter 30. Further, since this filter is of a film separating type, the secondary waste is not produced.

According to the system for collecting and refining SF₆ gas of the present invention, the deteriorated SF₆ gas is efficiently collected from the gas insulated machine, and impurities accompanying with the SF₆ gas are separated and removed, and the composition of the collected SF₆ gas can be measured and confirmed. Therefore, collecting, refining and reusing of the deteriorated SF₆ gas can be speedy performed at the installation site of the machine. Thereby, it is possible to prevent the SF₆ gas harmful to the global environment from releasing to atmosphere, and to make effective use of resources. Further, by using the dry type filter for the refining process, it is possible to make the system small in

size, to keep the portability of the system and to reduce the amount of the secondary waste.

Claims

- 5 1. A system for collecting and refining SF₆ gas comprising
 - a first filter (4) connected to a gas insulated machine (1),
 - a constant flow rate supplier (5) connected to the down-stream side of said first filter (4),
 - 10 a compressor (6) and a vacuum pump (7a) connected in parallel to the downstream side of said constant flow rate supplier (5),
 - a collecting tank (10) connected to the downstream side of said compressor (6), and
 - a cooler (20) for cooling said collecting tank (10),
 - wherein said first filter (4) is filled with a dry adsorbent.
- 15 2. The system of claim 1, comprising a constant flow rate valve arranged at the inlet side of said compressor (6).
3. The system of claim 1, wherein said first filter (4) is filled with a calcium group alkali.
4. The system of claim 1, comprising
 - 20 at least a second filter (8) arranged between said compressor (6) and said collecting tank (10), and
 - an analysis equipment (14) including a gaschromatograph at the downstream side of said second filter (8).
5. The system of claim 1, comprising a returning pipe for returning SF₆ gas from said collecting tank (10) to said gas
 - 25 insulated machine (1).
6. A method of collecting and refining SF₆ gas, comprising the steps of:
 - 30 removing impurities from SF₆ gas from a gas insulated machine (1) by averaging the SF₆ gas flow rate and letting the SF₆ gas flow through a first filter (4) filled with a dry adsorbent,
 - adsorbing and removing by-product gases by a second filter (8) filled with a dry adsorbent, and
 - collecting the SF₆ gas separated by said second filter (8) by cooling and liquefying the SF₆ gas.
7. The method of claim 6, comprising the further step of
 - 35 confirming compositions of said collected SF₆ gas by analysing compositions on analysis items of moisture concentration, acidic gas concentration, carbon dioxide concentration, any one of oxygen and nitrogen concentration and PFC concentration, and
 - 40 repeating said process of confirming said compositions and returning said collected SF₆ gas to said first filter (4) until said compositions become target values.

FIG. 1

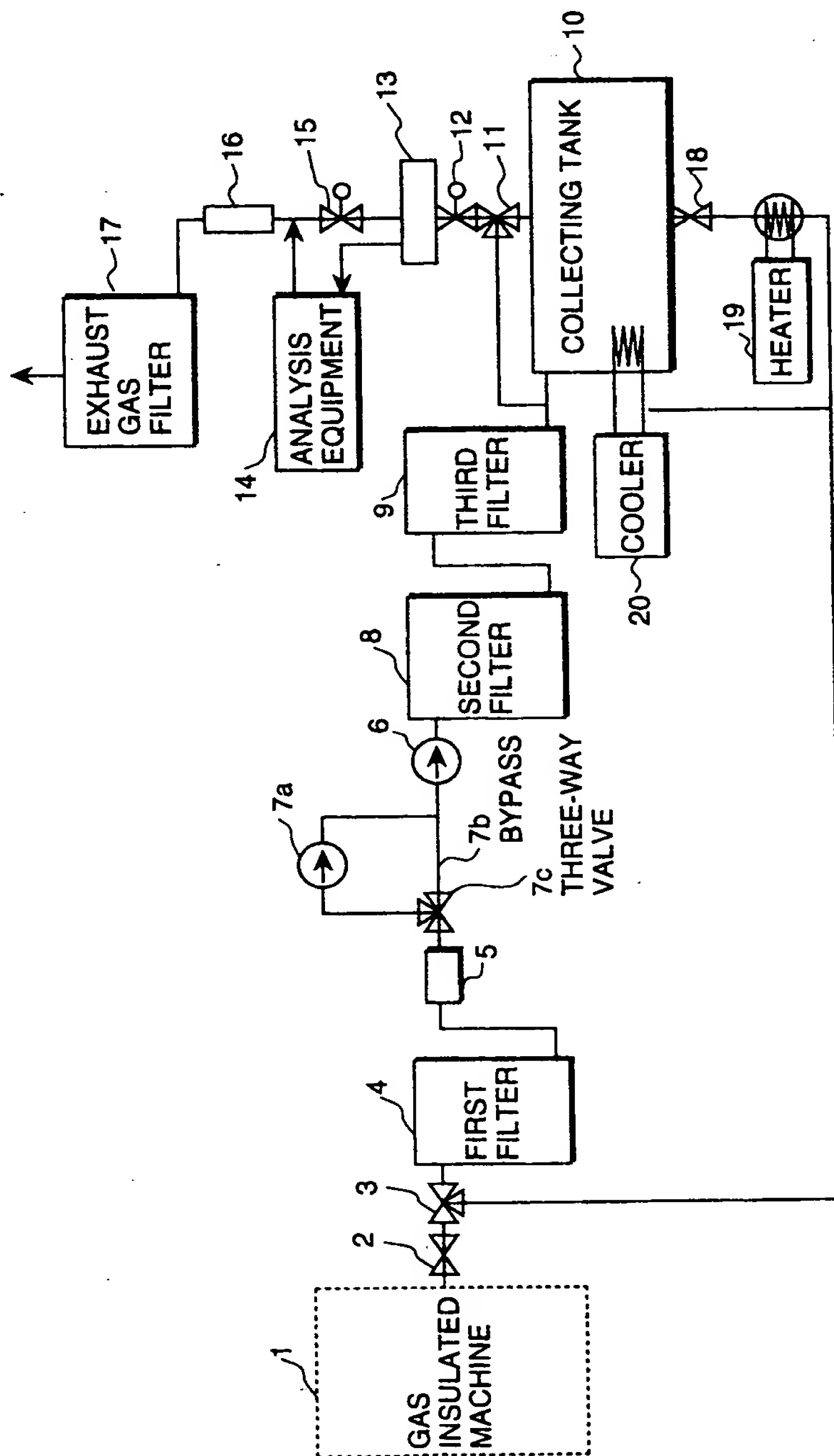


FIG.2

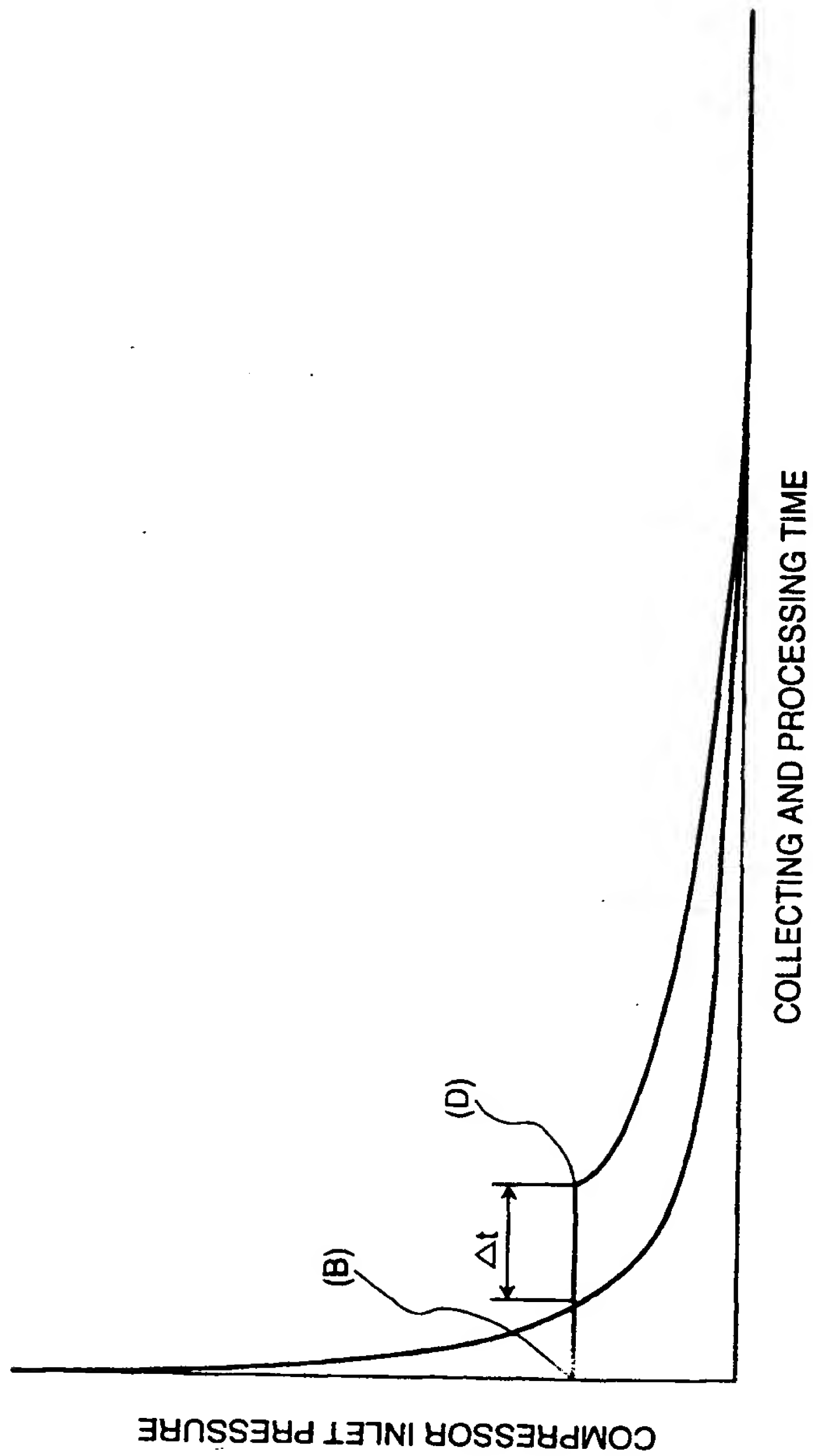


FIG.3

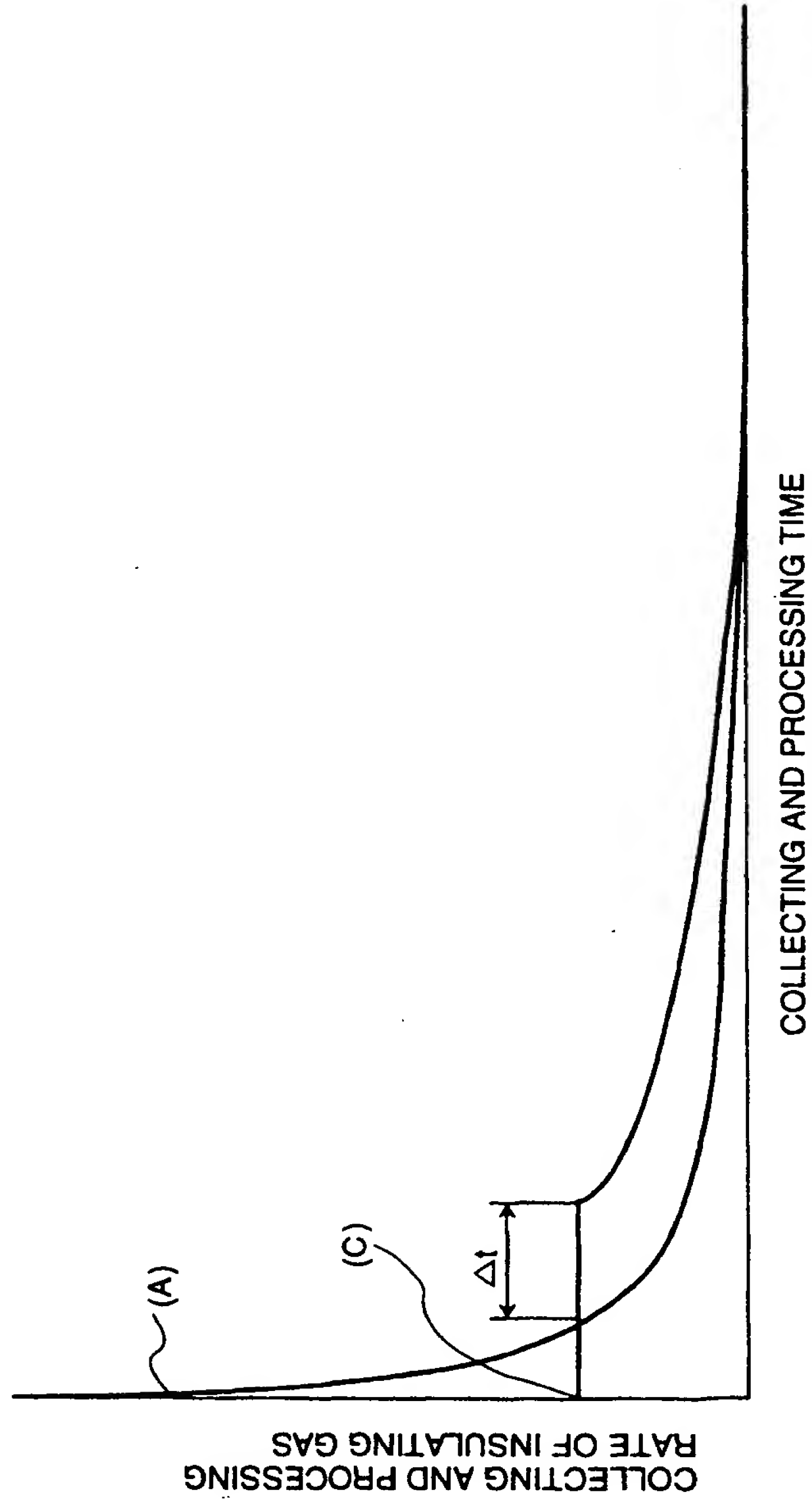


FIG. 4

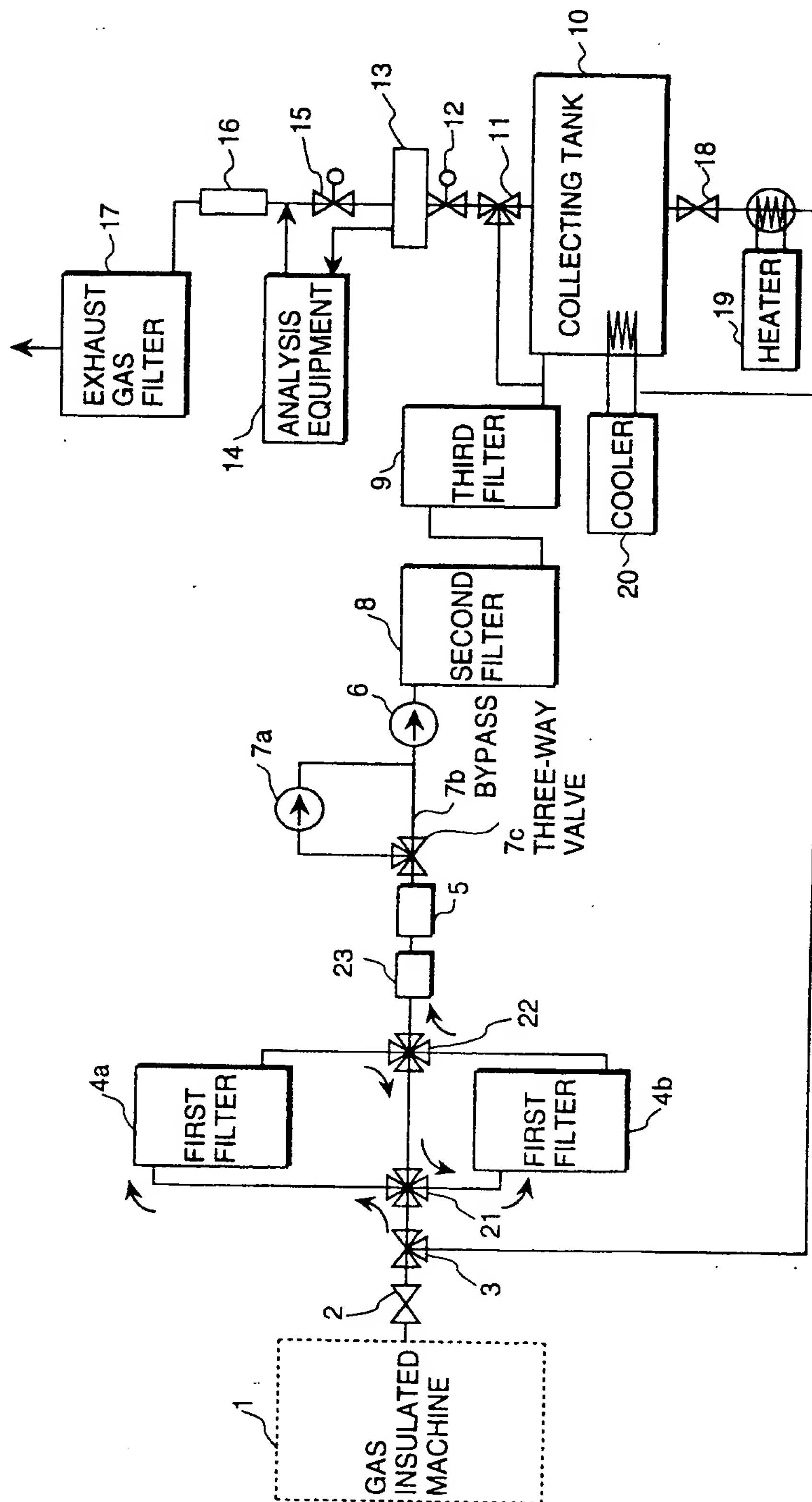


FIG. 5

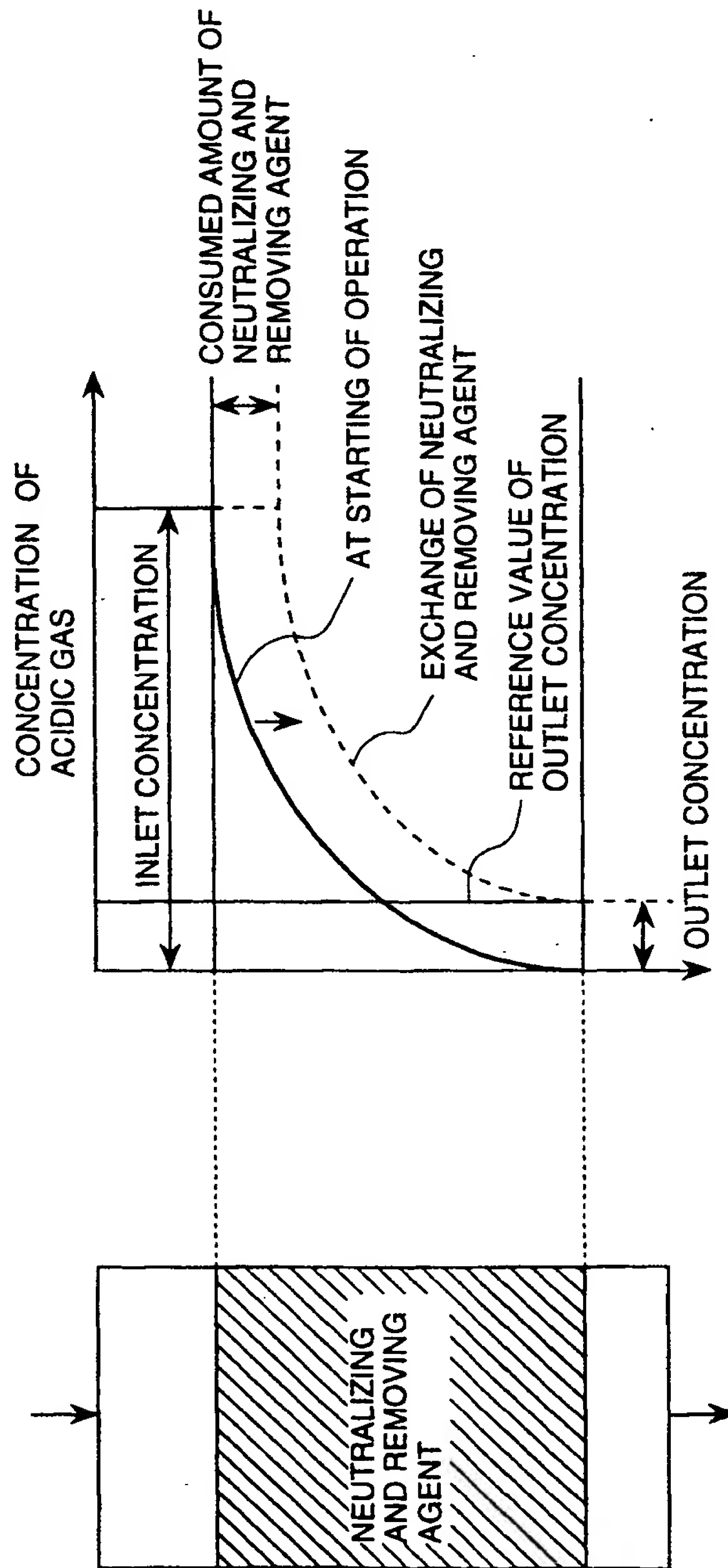


FIG. 6

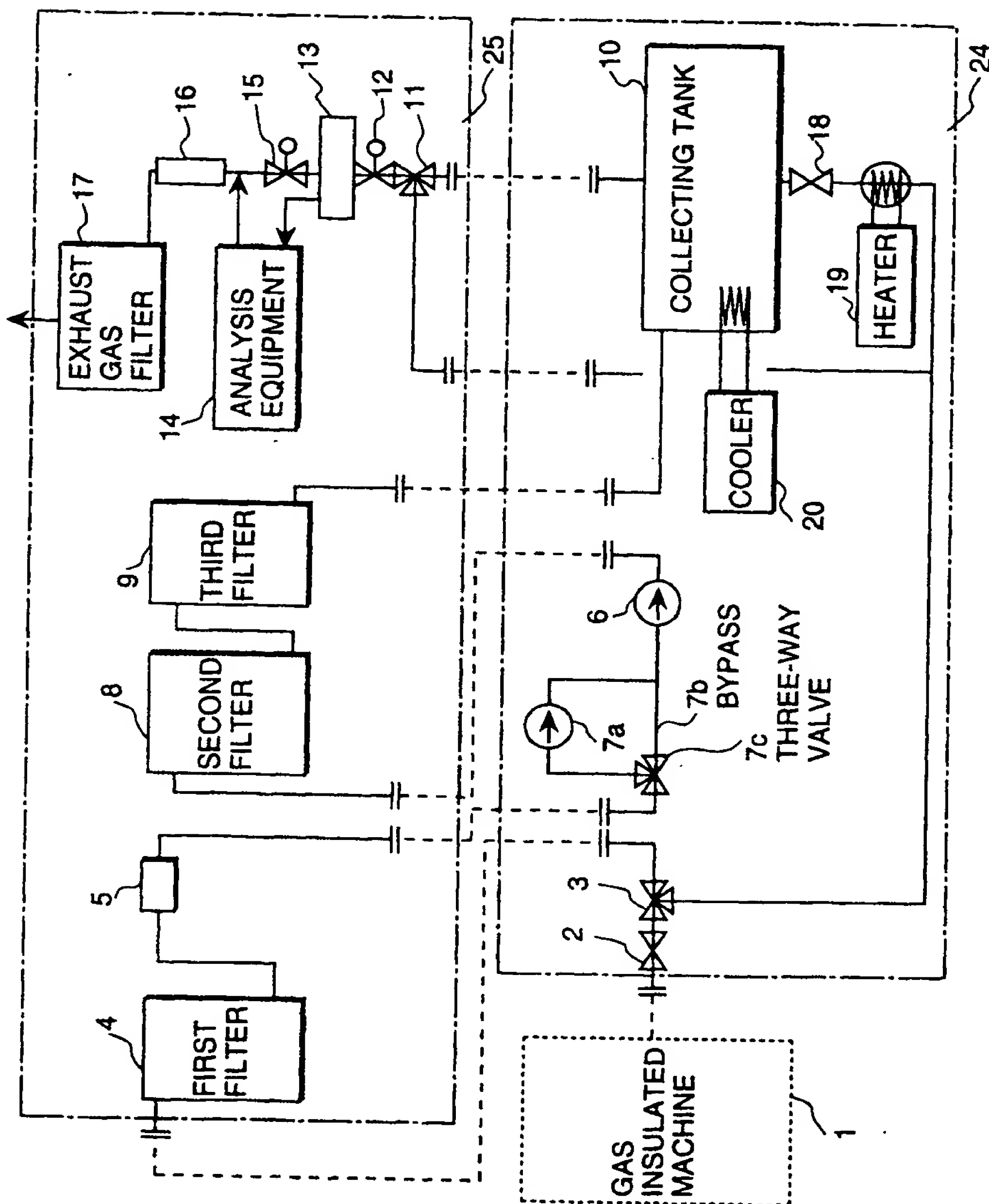


FIG. 7

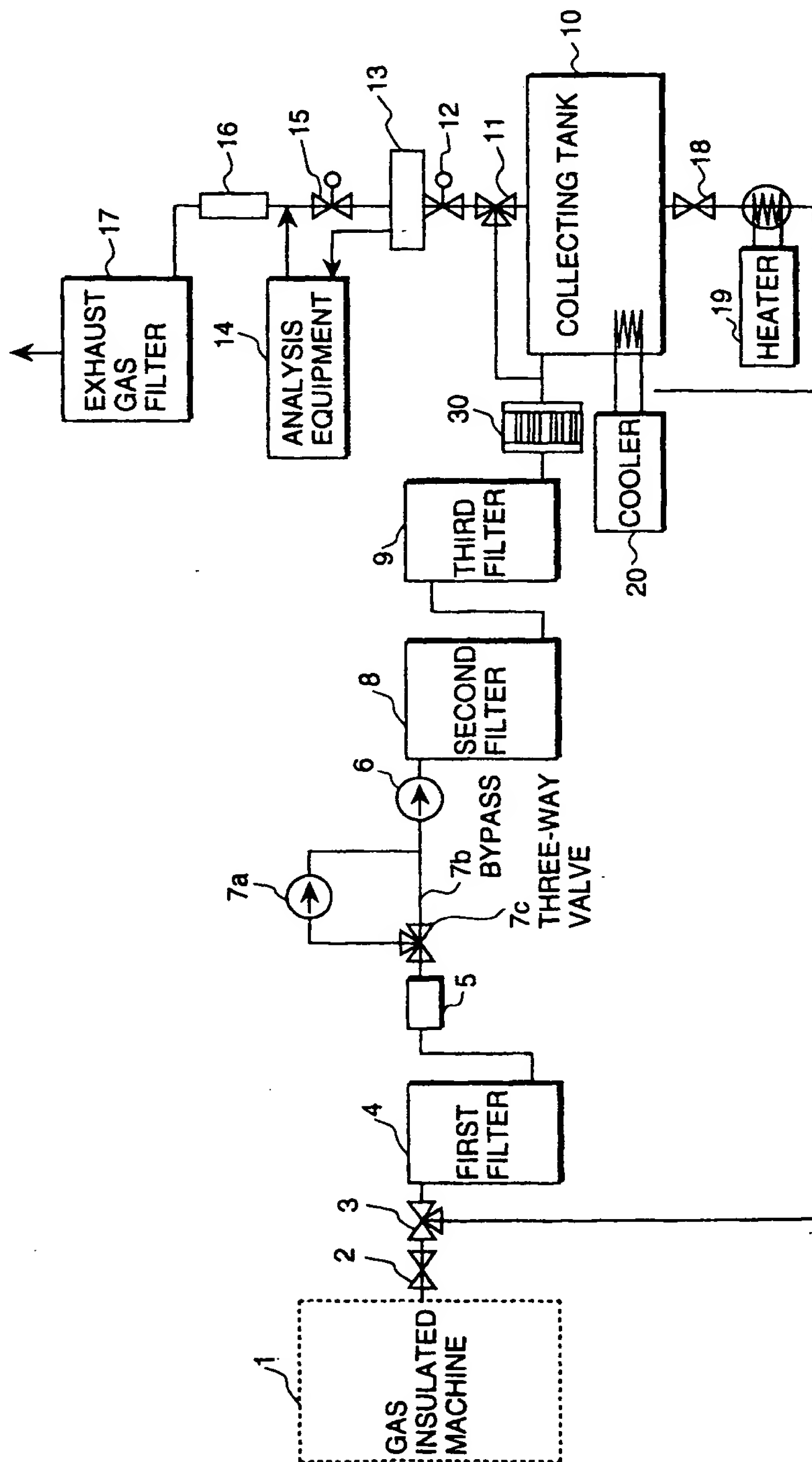
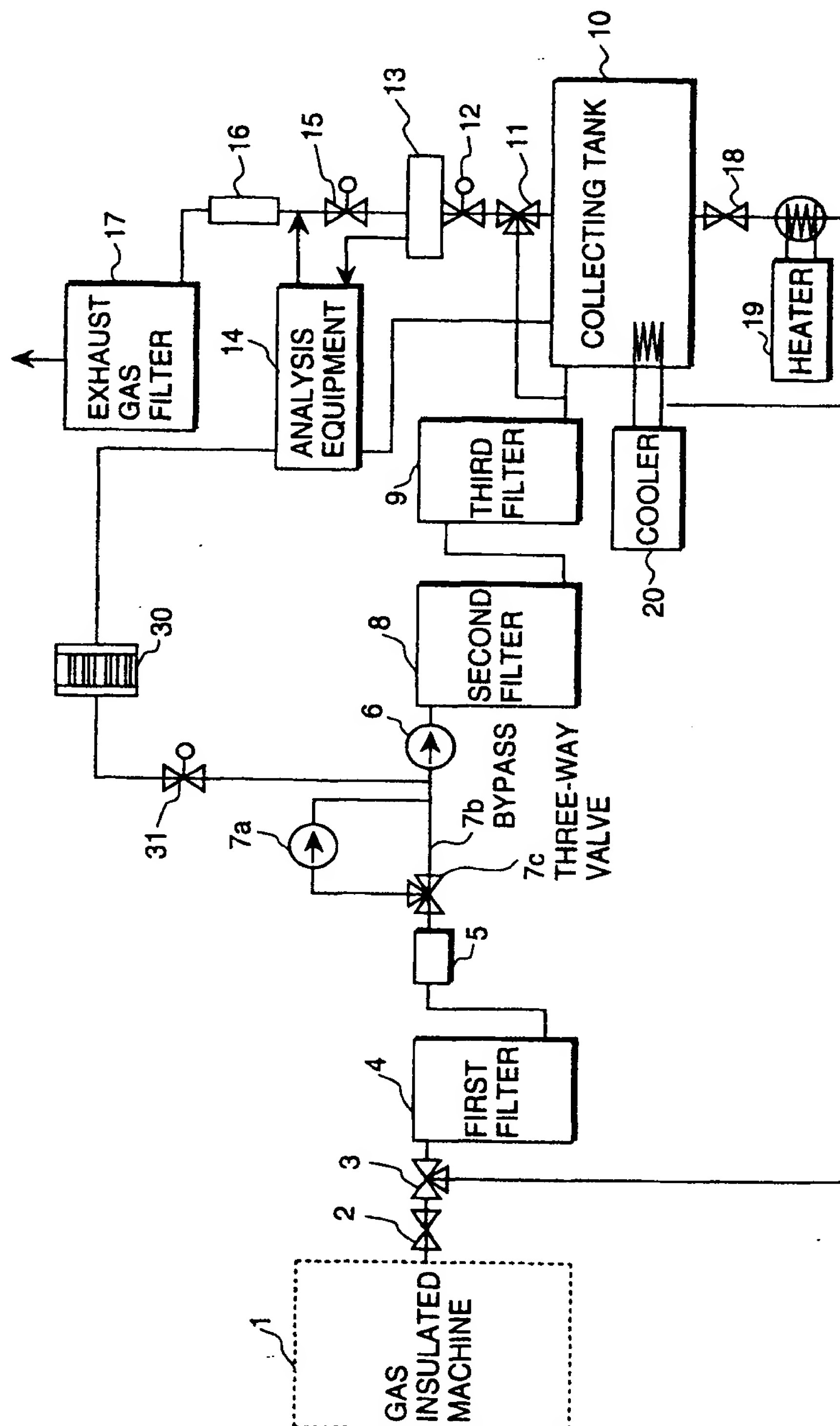


FIG. 8





European Patent
Office

EUROPEAN SEARCH REPORT

Application Number
EP 98 11 1045

DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
A	DATABASE WPI Section Ch, Week 9532 Derwent Publications Ltd., London, GB; Class A97, AN 95-241096 XP002077453 "Modified molecular sieve adsorbent for appts. - used for purifying sulphur hexafluoride gas" & CN 1 088 481 A (DALIAN CHEM PHYSICS INST CHINA ACAD SCI), 29 June 1994 * abstract *	1-5	C01B17/45 H02B13/055 B01D53/68 H01H33/56
A	DE 24 59 998 A (SIEMENS AG) 1 July 1976 * the whole document *	1-7	
A	EP 0 173 640 A (MITSUBISHI HEAVY IND LTD) 5 March 1986 * the whole document *	1-7	
A	GB 2 103 195 A (GNI ENERGET I IM GM KRZHIZHANO) 16 February 1983 * the whole document *	1-7	
A	WO 97 05635 A (FOERDERUNG DER SPEKTROCHEMIE U ;CENTRAL LAC CONVENIO COPEL E U (BR) 13 February 1997 * the whole document *	6,7	C01B H02B B01D H01H
The present search report has been drawn up for all claims			
Place of search THE HAGUE		Date of completion of the search 15 September 1998	Examiner Ross, R
CATEGORY OF CITED DOCUMENTS X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document			

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